

Systematic investigation of genetic vulnerabilities across cancer cell lines reveals lineage-specific dependencies in ovarian cancer

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A comprehensive understanding of the molecular vulnerabilities of every type of cancer will provide a powerful roadmap to guide therapeutic approaches. Efforts such as The Cancer Genome Atlas Project will identify genes with aberrant copy number, sequence, or expression in various cancer types, providing a survey of the genes that may have a causal role in cancer. A complementary approach is to perform systematic loss-of-function studies to identify essential genes in particular cancer cell types. We have begun a systematic effort, termed Project Achilles, aimed at identifying genetic vulnerabilities across large numbers of cancer cell lines. Here, we report the assessment of the essentiality of 11,194 genes in 102 human cancer cell lines. We show that the integration of these functional data with information derived from surveying cancer genomes pinpoints known and previously undescribed lineage-specific dependencies across a wide spectrum of cancers. In particular, we found 54 genes that are specifically essential for the proliferation and viability of ovarian cancer cells and also amplified in primary tumors or differentially overexpressed in ovarian cancer cell lines. One such gene, *PAX8*, is focally amplified in 16% of high-grade serous ovarian cancers and expressed at higher levels in ovarian tumors. Suppression of *PAX8* selectively induces apoptotic cell death of ovarian cancer cells. These results identify *PAX8* as an ovarian lineage-specific dependency. More generally, these observations demonstrate that the integration of genome-scale functional and structural studies provides an efficient path to identify dependencies of specific cancer types on particular genes and pathways.

high throughput | RNAi | oncogene | lineage

The application of whole-genome approaches to identify genetic alterations in cancer genomes is providing new insights into the spectrum of molecular events that occur in human tumors. Although in some cases this knowledge immediately illuminates a path toward clinical implementation, the long lists of genes with aberrant sequence, copy number, or expression in tumors already found make it clear that complementary information from systematic functional studies will be essential to obtain a comprehensive molecular understanding of cancer and to convert this knowledge into therapeutic strategies.

Most ovarian cancer patients present at an advanced stage with widely disseminated disease in the peritoneal cavity. Despite advances in surgery and chemotherapy, the majority of ovarian cancer patients re-present with relapsed and progressively chemotherapy-resistant and lethal disease. The Cancer Genome Atlas (TCGA) Project has characterized nearly 500 primary high-grade serous ovarian cancer genomes to identify recurrent genetic alterations in ovarian cancer (1).

A major feature of ovarian cancers is recurrent regions of copy-number alteration (1). A small number of these recurrent genomic events harbor known oncogenes and tumor suppressor genes, such as *MYC*, *CCNE1*, and *RB* (2). However, as in other cancers, the specific genes out of the ~1,800 genes targeted by recurrent amplification events remain undefined. Here, we report a genome-scale functional study to identify genes that are essential for the survival of 102 human cancer cell lines. The interrogation of a large number of cancer cell lines provides increased power to identify relationships between gene expression and dependence. To demonstrate the utility of integrating these data with results from cancer genome characterization, we focused on ovarian cancer and identified ovarian cancer lineage-specific dependencies.

Results

To identify genes essential for the proliferation and survival of human cancer cell lines, we performed genome-scale, pooled short hairpin RNA (shRNA) screens (3) in 102 cell lines (Fig. 1A and Fig. S1A), including 25 ovarian, 18 colon, 13 pancreatic, 9 esophageal, 8 non-small-cell lung cancer (NSCLC), and 6 glioblastoma cancer cell lines (Table S1). Each cell line was infected in quadruplicate (Fig. S2A) with a pool of lentivirally delivered shRNAs, composed of 54,020 shRNAs targeting 11,194 genes, and propagated for at least 16 doublings. The abundance of shRNA sequences at the endpoint relative to the initial reference pool was measured by microarray hybridization. We developed a standardized analytical pipeline to assess the effects on proliferation induced by each shRNA (Figs. S1B and S2B and C). Replicate infections were highly correlated (Fig. S2D and E), and hierarchical clustering demonstrated that the replicates clustered tightly together (Fig. 1B). In all, we obtained 22 million individual measurements of shRNA effects on proliferation.

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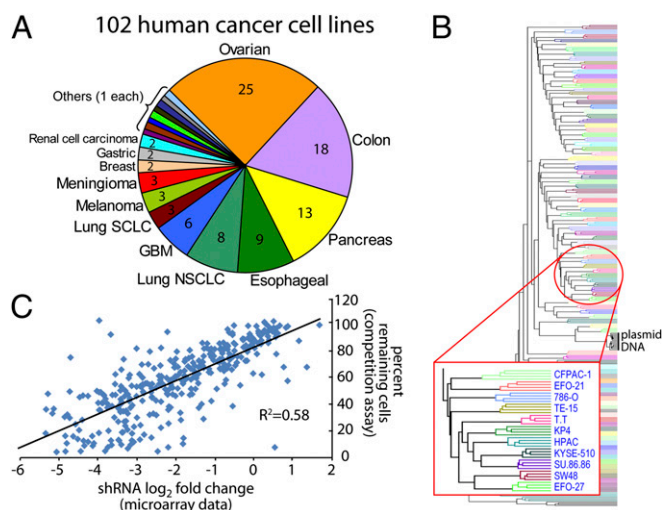


Fig. 1. Genome-scale RNAi screening identifies essential genes in 102 cancer cell lines. (A) Chart showing the number of cell lines from different lineages screened. (B) Unsupervised hierarchical clustering of the shRNA hybridization data obtained from quadruplicate screens of 102 cancer cell lines (various colors) and the shRNA plasmid DNA reference pool (15 replicates). A representative portion of the dendrogram is depicted at higher magnification. (C) The relative abundance 7 d after infection of OVCAR-8 cells infected with 350 individual shRNAs encoded in a GFP+ plasmid (y axis) correlates with the relative abundance (\log_2 fold change) of each shRNA measured in the pooled shRNA screen by microarray hybridization (x axis).

To ensure that the pooled screening process accurately measured the activity of the shRNAs, we individually retested 350 shRNAs, chosen to sample the full range of array measurements obtained in a pooled screen in OVCAR-8 cells, by performing competition assays (Fig. 1C and Fig. S3). Specifically, we cloned these shRNAs into a modified pLKO.1 viral vector that coexpresses GFP and used these vectors to introduce shRNAs into ~50% of an OVCAR-8 cell population. We then monitored the proportion of GFP-expressing cells over time to measure the effects of each shRNA on proliferation. The percent depletion of the shRNAs in the individual pairwise competition tests was correlated to the $\log(\text{fold depletion})$ of these shRNA in the pooled screen ($R^2 = 0.58$; Fig. 1C and Table S2). We note that shRNAs that show the largest degree of depletion in the pooled screen exhibited more variability as would be expected at the limit of signal detection. These two correlated assessments were made at different time points, further confirming that they provide robust measures of the intrinsic proliferation effects of the individual shRNAs.

Correlation of Genetic Dependency with Properties of Cell Lines. We then sought to understand how the vulnerabilities of cancer cell lines relate to various properties, such as mutations in a specific gene, disruption of a specific pathway, or inclusion in a specific lineage. Because human cancer cell lines are genetically and epigenetically diverse, the analysis of a large number of cell lines ensures that the relationships between cell properties and the dependence of those cells on specific genes are not particular to one context. We tested whether this large dataset permits reliable inferences about the genetic vulnerability of cancers possessing specific properties. For each cell line classification, we used a class-discrimination feature selection method called the “weight of evidence” (WoE) statistic (4, 5) to rank shRNAs by their ability to distinguish the specified classes.

Dependencies of Cell Lines with Oncogenic Mutations. We first examined vulnerabilities of cell lines with *KRAS* or *BRAF* mutations. We defined “essential” genes by three complementary methods, including (i) WoE rank of the top shRNA targeting

each gene, (ii) WoE rank of the top two shRNAs targeting each gene, or (iii) a composite score of WoE ranks for all shRNAs for each gene using the Kolmogorov–Smirnov (KS) statistic (3). The *KRAS* and *BRAF* genes themselves were ranked highly by all three methods (Fig. 2A and D). The top scoring *KRAS*- and *BRAF*-specific shRNAs significantly discriminated between the mutant and wild-type classes ($P = 1.89 \times 10^{-5}$ and 1.89×10^{-4} , respectively, WoE; Fig. 2B and E).

Interestingly, because 7/10 *BRAF*-mutant cell lines were derived from nonmelanoma lineages (including 5 from colon cancer), these observations suggest that cancer cell lines that harbor mutant *BRAF* exhibit a similar dependence on *BRAF*. Although initial reports suggest that clinical responses to *BRAF* inhibition in *BRAF*-mutant colon tumors are much less robust than those observed in *BRAF*-mutant melanomas (6), our observations indicate that *BRAF* is essential in colon cancer cell lines that express mutant *BRAF* (Fig. S4A and B).

We next examined vulnerabilities in cell lines harboring *PIK3CA* mutations. *PIK3CA* itself strongly scored as a top differentially essential gene between *PIK3CA* mutant and *PIK3CA* wild-type cell lines in 2/3 gene-level analyses (Fig. 2G and H). *MTOR* ranked highly in 2/3 analyses (23rd and 30th of 11,194; Fig. S4C); the top scoring *MTOR*-specific shRNA strongly discriminated the *PIK3CA* mutant and wild-type classes ($P = 6.03 \times 10^{-4}$, WoE; Fig. S4D), indicating that cell lines that harbor *PIK3CA* mutations are also dependent on mTOR. These observations confirm prior work showing that mTOR plays an important role in PI3K signaling (7).

To assess how the number of cell lines analyzed affected these analyses, we repeated our WoE scoring of a set of shRNAs using data from smaller numbers of cell lines subsampled from the entire dataset. For the top scoring *KRAS*, *BRAF*, and *PIK3CA* shRNAs that were able to distinguish cell lines with mutant or wild-type alleles of these same respective genes, we found that the analysis of a smaller number of cell lines (<5) decreased the likelihood of discriminating between these two classes, whereas the comparison of groups composed of >10 cell lines greatly increased our ability to distinguish the two classes (Fig. 2C, F, and I). We concluded that the analysis of a large number of cell lines overcomes the inherent heterogeneity of cell lines and reveals robust relationships between essentiality and particular cell features that persist across different genetic and epigenetic backgrounds. With this foundation, we undertook a preliminary exploration of what can be learned about genetic vulnerabilities of cancer cells with specific properties.

Lineage-Specific Genetic Dependencies. We hypothesized that a subset of genes showing enhanced dependencies in specific lineages would also be aberrantly activated in tumors due to amplification or overexpression. Recent studies have identified oncogenic transcription factors that are amplified, overexpressed, and essential in subsets of tumors from cancers of specific lineages, including *NKX2-1* in lung adenocarcinoma (8), *MITF* in melanoma (9), and *SOX2* in squamous cell carcinomas (10). To identify lineage-specific dependencies, we ranked shRNAs by their ability to discriminate cell lines of one lineage from cell lines from all other lineages (Fig. 3A). We selected the top 150 genes (1.3% of those screened) based on ranking of the top-ranked shRNAs, the top 300 genes (2.7%) by the second-best ranking shRNAs, and the top 300 genes (2.7%) as assessed by the KS statistic (ref. 3; Table S3). Three categories of essential genes were considered for further analysis: (i) genes scoring by all three methods from individual lineage analyses; (ii) genes scoring by any method that also were amplified in primary tumors (essential and amplified; Table S3); and (iii) genes scoring by any method that also were differentially up-regulated in cell lines derived from that lineage (essential and overexpressed; Tables S3 and S4). An overview of these results from analyses performed across six cancer lineages is displayed in Table S5.

In colon cancer cell lines, we found *KRAS*, *CTNNB1*, and *BRAF* among 23 essential genes that scored by all three methods and

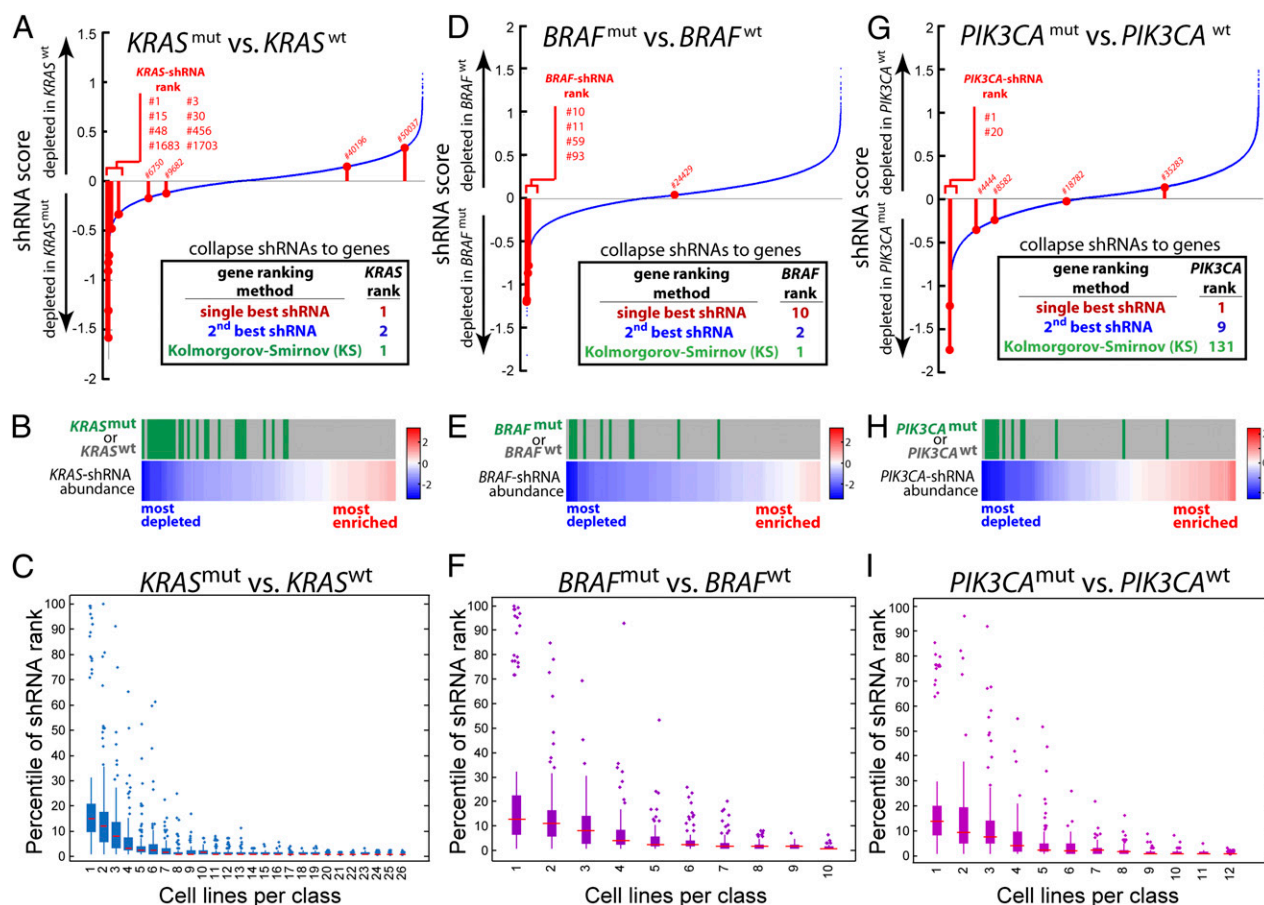


Fig. 2. Dependencies of cell lines with mutations in *KRAS*, *BRAF*, or *PIK3CA*. (A, D, and G) Distribution of shRNA ranks (x axis) by the WoE scores (y axis) for the class comparisons of *KRAS* mutant (mut) vs. *KRAS* wild-type (wt) cell lines (A), *BRAF* mutant vs. *BRAF* wt (D), and *PIK3CA* mutant vs. *PIK3CA* wt (G). shRNAs targeting *KRAS*, *BRAF*, and *PIK3CA* are highlighted in red, and their ranks are listed. Insets report the gene ranks of *KRAS*, *BRAF*, and *PIK3CA* for differential essentiality in the subset of cell lines with activating mutations in those respective genes. (B, E, and H) *KRAS* (B), *BRAF* (E), or *PIK3CA* (H) mutation status (mutant lines in green, wt lines in gray) correlates strongly with depletion of shRNAs targeting these genes. (Lower) Heatmaps report *KRAS*- (B), *BRAF*- (E), and *PIK3CA*-shRNA (H) fold depletion in each cell line. (C, F, and I) Subsets of the 102 cell lines were analyzed to assess convergence of the gene dependency results for the *KRAS* (C), *BRAF* (F), and *PIK3CA* (I) mutant vs. wt class comparison analyses as a function of the number of cell lines tested. Distributions of the scores of the top *KRAS*, *BRAF*, and *PIK3CA* hit shRNAs (given as the percentile of their depletion rankings, with a smaller percentage corresponding to more depleted, y axis) in the respective cell line class comparisons (using WoE) are shown for each of 100 trials, subsampling the indicated numbers of cell lines in each class (mutant and wt). The red bar indicates the median value for each group of subsamplings, boxes represent the 25th to 75th percentile of the data, and whiskers extend to the most extreme values of the group that are not considered outliers.

found *KRAS* and *IGF1R* (11) among 35 essential and amplified genes. In pancreatic cancer cell lines, we identified *KRAS* among 23 essential genes that scored in all three analyses (Table S5). In NSCLC, we found *NKX2-1* as the only essential gene that is both amplified and overexpressed and found *CDK6* among 7 essential and amplified genes (8). These observations provide evidence that such integrative lineage analyses identify both known oncogenes and other relevant lineage-restricted dependencies.

In addition, we identified a number of particularly interesting, previously undescribed candidate lineage-specific dependencies. For example, we found *MAP2K4*, an activator of JNK and p38 (12), among 10 genes that showed selective essentiality and expression in NSCLC. We found *MYB* (13) and *AXIN2* (14) among 9 genes that were essential and differentially expressed in colon cancer, and we identified *SOX9* (15) among 18 genes in pancreatic cancer that emerged as lineage-specific dependencies nominated by all three gene-scoring methods.

To extend these observations, we selected the ovarian lineage for deeper analysis. Of the 582 genes (5.2%) nominated as candidates for enhanced dependency in ovarian cancer cells, we identified 22 essential genes that scored in all three analytical methods (Fig. 3B and Table S5) and found 5 essential and

overexpressed genes (Tables S4 and S5). TCGA identified 1,825 genes residing on recurrently amplified regions in ovarian tumors, and we identified 50 amplified genes as also essential (Fig. 3B and Table S5). The set of amplified and essential genes included the known oncogene *CCNE1* (16) and candidates including the *FRS2* adaptor protein (17), the *PRKCE* protein kinase (18), *RPTOR* (19), and the *PAX8* paired box transcription factor. Similarly to *NKX2-1* in NSCLC, *MITF* in melanoma, and *MYB* in colon cancer, *PAX8* was not only essential and amplified but also overexpressed in a lineage-specific manner.

Characterization of *PAX8* Dependency in Ovarian Cancer. *PAX8* was the only gene that was (i) identified as an essential gene in all three scoring methods, (ii) amplified in primary high-grade serous ovarian tumors, and (iii) differentially expressed in ovarian cancer cell lines (Table S5). Cell line subsampling analysis revealed that the large number of ovarian cell lines screened enabled the identification of this previously undescribed dependency (Fig. 3C).

PAX8 is a lineage-restricted transcription factor that plays an essential role in organogenesis of the Müllerian system (20), the thyroid, and the kidney (21). *PAX8* was previously found to be

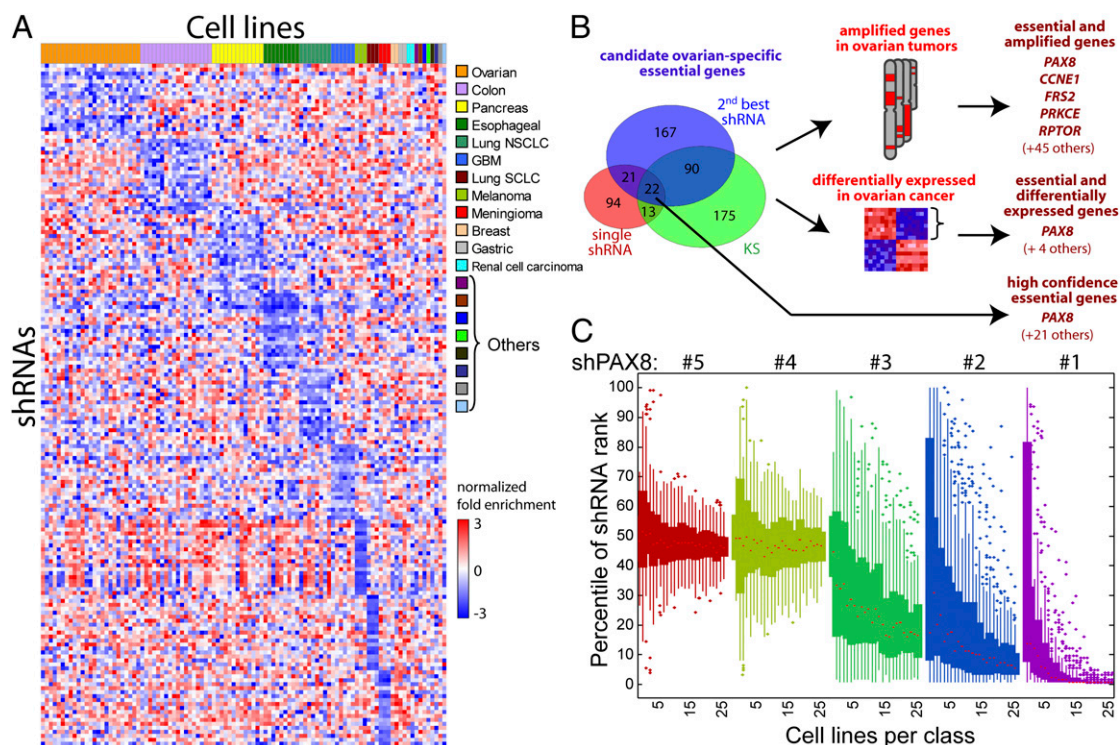


Fig. 3. Lineage-specific dependencies. (A) Heatmap of differentially antiproliferative shRNAs in cell lines from individual cancer lineages in comparison with all others. The top 20 shRNAs that distinguish each lineage from the others are displayed. (B) Ovarian-specific dependencies. Three complementary methods of gene scoring [ranking by (i) best or (ii) second best scoring shRNA or (iii) composite of all shRNAs for the gene using a KS statistic identified 582 (5.2%) genes that were selectively required for ovarian cancer cell proliferation. Fifty of these were among the 1,825 recurrently amplified genes in primary high-grade serous ovarian tumors (1). Among the 200 genes that were differentially overexpressed in ovarian cancer cell lines, 114 genes were included in the shRNA pool, and 5 genes showed enhanced essentiality in ovarian cancer lines. Twenty-two of these genes that were scored by all three gene-scoring methods were considered as high-confidence essential genes. (C) Distributions of the scores of PAX8 shRNA (given as the percentile of their rankings, y axis) after 100 trials of the ovarian vs. nonovarian WoE comparison, for equal class sizes of 1–25 (x axis; colors indicate PAX8 shRNAs 1–5). The red bar indicates the median value for each group of subsamplings, boxes represent the 25th to 75th percentile of the data, and whiskers extend to the most extreme values of the group that are not considered outliers.

overexpressed in ovarian cancers (22) and implicated in follicular thyroid cancer development (23). We observed that *PAX8* was the most differentially expressed gene when we compared ovarian cell lines to all of the other cancer cell lines (Fig. 4A). Furthermore, we found that *PAX8* was amplified in 16% of primary ovarian tumors [$\log_2(\text{copy number ratio}) > 0.3$; $n = 345$] in a peak (2q13) that also contains *PSD4* and *LOC654433* (Fig. 4B).

cells (Fig. 4D). Suppression of *PAX8* induced apoptosis in these ovarian cancer cell lines (Fig. 4F). In contrast, the two cell lines (COV504 and OV-90) least sensitive to *PAX8* suppression did not harbor the 2q13 amplification and expressed relatively low levels of *PAX8* (Fig. 4D). These observations suggest that *PAX8* represents a lineage-specific essential gene in a significant subset of ovarian cancer.

Discussion

The integrated analysis of functional dependencies and alterations in cancer genomes presented herein identified potential targets in ovarian, lung, colon, glioblastoma, pancreatic, and esophageal cancers. Among these candidate genes, we identified known oncogenes and lineage-specific dependencies as well as previously undescribed candidates, including the *PAX8* transcription factor in ovarian cancer. Although shRNA screens performed in small numbers of cell lines have identified essential genes in specific contexts, the interrogation of genes across a large number of human cancer cell lines through Project Achilles provides a substantially more robust assessment of gene dependence and overcomes confounding effects due to the inherent heterogeneity of human cancer cell lines. These datasets will enable a wide range of analyses to connect particular cancer genotypes to dependencies.

As an initial approach, we elected to explore dependencies harbored by a majority of ovarian cancers. We pinpointed 5 genes displaying enhanced essentiality in ovarian cancer and differential overexpression in ovarian cell lines and 50 genes displaying enhanced essentiality in ovarian cancer and amplification in ovarian tumors. Further studies will be necessary to

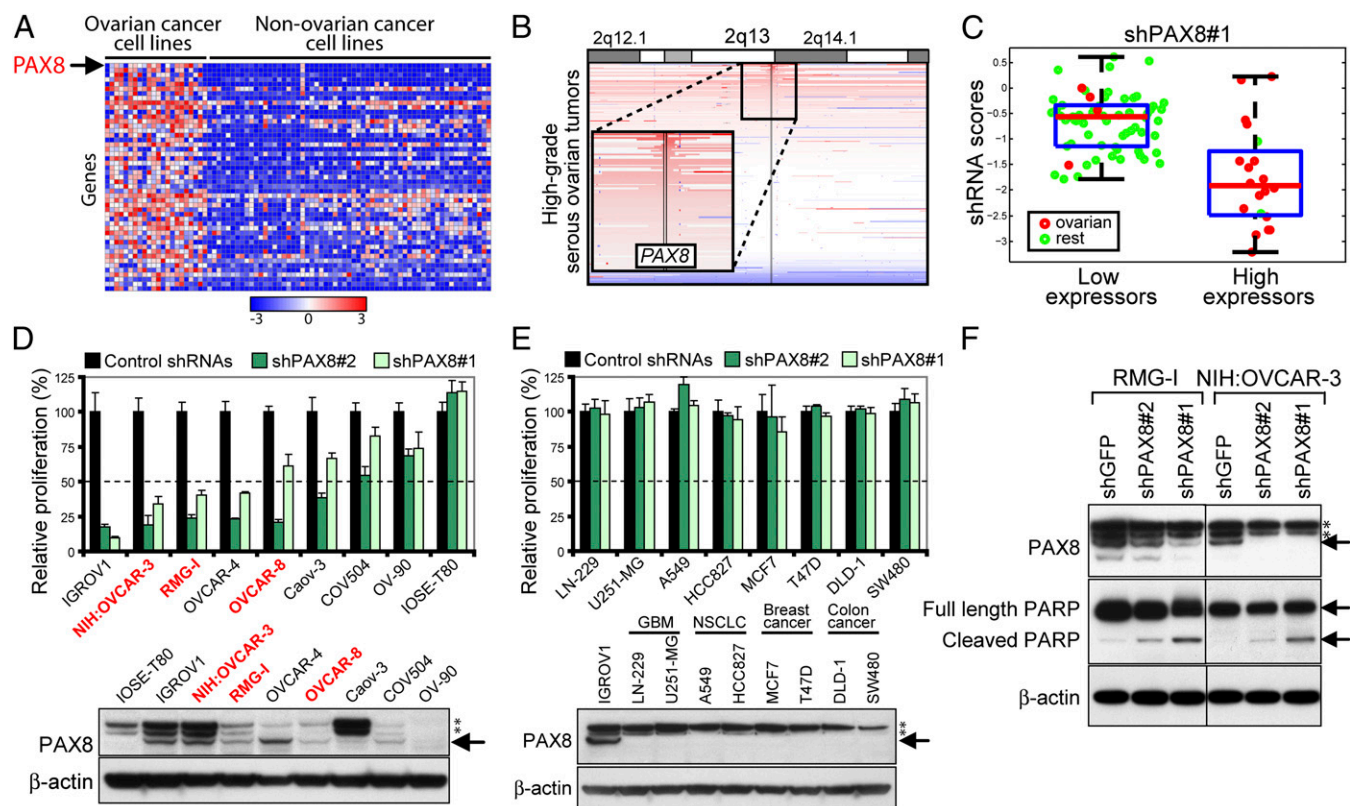


Fig. 4. *PAX8* is essential for ovarian cancer cell proliferation and survival. (A) *PAX8* is the top-ranked differentially expressed gene between ovarian and nonovarian cancer cell lines. Arrow indicates *PAX8*. (B) SNP array colorgrams depict genomic amplification of *PAX8* in primary high-grade serous ovarian cancers (1). Regions of genomic amplification and deletion are denoted in red and blue, respectively. SNP array profiles derived from primary ovarian tumors were sorted based on the degree of amplification of each gene. Black vertical lines denote the boundaries of the *PAX8* gene. (C) Boxplot showing significant difference in the degree of depletion of a *PAX8*-specific shRNA in 63 cell lines with low levels of *PAX8* compared with 20 lines with high levels of *PAX8* ($P = 2.14 \times 10^{-8}$, t test). Cell lines were divided into high- and low-expressing groups. The red line indicates the median value for each group, boxes represent the 25th to 75th percentile of the data, and whiskers extend to the most extreme values of the group that are not considered outliers. Ovarian cancer cell lines are plotted with red circles; cell lines from all other lineages are plotted with green circles. (D Upper) Effects of *PAX8* suppression on proliferation in eight ovarian cancer cell lines; dotted line indicates 50% relative proliferation. (Lower) Immunoblot of *PAX8* in a panel of eight ovarian cancer cell lines and in immortalized IOSE-T80 cells. Cell lines with amplification of 2q13 (\log_2 copy number ratio > 0.3) are marked in red. * denotes nonspecific band. (E Upper) Effects of *PAX8* suppression on proliferation of cell lines from indicated cancer types. (Lower) Immunoblot of *PAX8*. Error bars indicate SD of six replicate measurements. (F) Immunoblot of poly(ADP-ribose) polymerase after *PAX8* suppression in two 2q13-amplified cell lines. * denotes nonspecific band.

extend the preliminary findings for these ovarian cancer dependencies. A single gene, *PAX8*, emerged from every one of these analyses. Our subsequent experiments confirmed *PAX8* to be a lineage-specific survival gene that is essential for proliferation of ovarian cancer cells, highly expressed in ovarian cancer lines, and amplified in a substantial fraction of primary ovarian tumors.

PAX8 has been shown to play an essential role in the normal development of the thyroid gland (21) and female genital tract (20). The thyroid gland in *Pax8*-deficient mice is completely devoid of thyroid hormone-producing follicular cells and exhibits severe growth retardation within a week after birth (21). Thyroxine substitution enables *Pax8*-deficient mice to survive to adulthood (20). However, these mice are infertile because of the absence of uterus and vaginal openings (20), indicating that *Pax8* is also essential for the development of the Müllerian duct. In the reproductive tract, *PAX8* expression is restricted to secretory cells of the fallopian tube epithelium (22), which recent reports suggest represent a cell of origin for serous ovarian cancer (25). These findings suggest that *PAX8* plays critical roles both in normal development of female genital tract and in high-grade serous ovarian tumors.

A number of genes involved in the differentiation programs for specific tissue lineages are amplified in cancers that arise from these tissues. For example, *NKX2-1* (8), *MITF* (9), and *SOX2* (10)

are amplified and essential for the survival of significant subsets of NSCLC, melanoma, and squamous cell carcinomas, respectively. Our results show that *PAX8* belongs to this distinct class of lineage-survival genes that are required for both normal development of specific tissues and for cancer cell proliferation/survival. Although different subtypes of ovarian cancer are likely to harbor distinct profiles of genetic alterations (2), our findings suggest that for most ovarian cancer cell lines, *PAX8*-driven transcription programs transiently active during normal development are coopted to maintain the malignant state.

Epithelial ovarian cancers have been classified into four major subtypes based on histology: serous, clear cell, endometrioid, and mucinous (2). These major subtypes show morphologic features that resemble those of the epithelia of the reproductive tract derived from the Müllerian duct (2). Previous studies using immunohistochemistry showed that 89–100% of serous, clear cell, and endometrioid subtypes and 8% of mucinous subtype express *PAX8* (26). The majority of the ovarian cancer cell lines that we screened (20/25) belong to the serous subtype or had a mixed morphology. A larger collection of cancer cell lines will facilitate a deeper investigation of ovarian cancer subtypes.

Given the genetic heterogeneity of cancer, screening large numbers of cell lines is required to have sufficient statistical power to extract known and novel relationships and provide adequate representation of individual sublineage classes. Although the

classification of cell lines based on a single characteristic is unlikely to segregate cell lines into homogenous groups, our observations indicate that by including many cell line representatives of each class of interest, it is still possible to discover underlying relationships between genotype or lineage and essential genes. Integration of these data with information from the analyses of structural alterations in cancer genomes will further facilitate the systematic identification of genes critical to oncogenesis.

The approach described here can be extended to enable systematic interrogation of codependences beyond those analyzed here, including synthetic lethal relationships with activated oncogenes or inactivated tumor suppressor genes (27). Analyses of dependencies of cell lines with particular characteristics have the potential to discover novel targets, for example, by integrating the output of these types of screens with information emerging from the cataloging of mutations and other alterations in cancer genomes.

More generally, as large-scale efforts to characterize cancer genomes accelerate, these observations illustrate a path to functionally characterize the genes found to be altered in tumors and to identify the subset of such genes critical to cancer initiation and maintenance. To this end, we have made this dataset available (www.broadinstitute.org/igp) and will update the Project Achilles database as more data are obtained. Beyond the specific findings reported herein, we anticipate that this dataset will prove useful to identify correlations between genetic features and essential genes in human cancer cell lines.

Materials and Methods

Pooled shRNA Screening. Lentiviral pLKO.1- shRNA constructs were obtained from the RNAi Consortium, and the human 54K pool of 54,020 shRNA plasmids was assembled by combining 16 normalized subpools of ~3400 shRNA plasmids. The list of 54,020 shRNAs can be found at <http://www.broadinstitute.org/igp>. Genome-scale pooled shRNA screens to identify genes essential for proliferation in 102 cancer cell lines were performed (3) using a lentivirally delivered pool of 54,020 shRNAs targeting 11,194 genes. The culture conditions for all cancer cell lines are listed in Table S1. Each cell line was infected in quadruplicate and propagated for at least 16 population doublings. The abundance of shRNA constructs relative to the initial DNA plasmid pool was measured by microarray hybridization (3) and analyzed by

using a uniform pipeline. Detailed descriptions of each procedure can be found in *SI Methods*.

Data Processing, Class Comparison, and Gene Ranking. Raw .CEL files from custom Affymetrix barcode arrays were processed with a modified version of dCHIP software (3). The GenePattern modules shRNAscores and NormalizeCellLines were used to calculate the log fold change in shRNA abundances for each cell line at the conclusion of the screening relative to the initial plasmid DNA reference pool and to normalize these depletion values by using peak median absolute deviation normalization, a variation of Z score with median absolute deviation (3). Class definition files (.cls) were made by using the GenePattern module SubsetGctandCls; definitions included cell line lineage (e.g., ovarian cancer, NSCLC, etc.) or genetic alterations (28, 29). To compute the statistical evidence that a given shRNA contributes to the observed essentiality phenotype between two classes of interest, we used a WoE approach (4, 5). The GENE-E program (<http://www.broadinstitute.org/cancer/software/GENE-E>; ref. 3) was used to collapse shRNAscores to gene rankings by three complementary methods. These methods included (i) ranking genes by their highest shRNA depletion score, (ii) ranking genes based on the P value rank (correcting for different set sizes of shRNA targeting different genes) of their second best ranked shRNA, and (iii) ranking genes using a KS statistic in an approach similar to gene set enrichment analysis (RNAi gene enrichment ranking; ref. 3). Detailed descriptions of each procedure can be found in *SI Methods*. All data files, accessory files, and GenePattern modules can be found through the Integrative Genomics Portal (<http://www.broadinstitute.org/igp>).

Competition Assay. OVCAR-8 (5×10^4) cells were seeded into each well of a 96-well plate and spin-infected with 2 or 4 μ L of lentiviruses (in duplicate) at $930 \times g$ for 2 h at 30 °C in the presence of 4 μ g/mL polybrene to transduce ~50% of the cells. Cells were then trypsinized and replated into 24-well plates. The percent of GFP+ cells at 3 and 7 d after infection was measured using BD LSR II flow cytometry system equipped with a high-throughput sampler (BD Biosciences). The fraction of GFP+ cells 7 d after infection relative to 3 d after infection was calculated. Data represent mean \pm SD of duplicate infections.

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